The 3_{1/3} Bands of Carbon Disulfide and Carbon Dioxide

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The $3\nu_3$ bands of carbon disulfide and carbon dioxide are described. This band of CS_2 is complex, containing two transitions from excited states and two transitions attributed to the isotopic molecule $CS^{32}S^{34}$ in addition to the primary transition $000\rightarrow003$ of CS_2^{32} . A multiple-reflection cell of 6-meter path length, 20-cm optics, and negligible loss of aperture is described. Molecular constants are checked for CO_2 and found for CS_2 . These latter constants are $B_0 = 0.1089$ cm⁻¹ ± 0.0004 , $\alpha_3 = 0.0007$ cm⁻¹ ± 0.00003 , $X_{33} = -5.2$ cm⁻¹, and $X_{23} = -3.3 \text{ cm}^{-1}$.

The infrared absorption spectrum of carbon dioxide is well known. Carbon disulfide, however, although of similar structure, has received much less study. This is probably due to the difficulty of resolving the fine structure of its spectrum, caused by the large moment of inertia of this molecule. In order to resolve the fine structure, several requirements must be met. A spectrometer capable of resolving to 1 or 2 tenths of a wave number in the region of the spectrum must be used, and a path of sufficient length to permit measurable absorption must be devised.

Both conditions being obtainable in the Radiometry laboratory of the Bureau, the $3\nu_3$ band of CS₂ has been studied, along with the analogous band of CO₂. This latter band has been observed under low resolution by Barker and Wu¹ and under higher resolution by Goldberg, Mohler, Pierce, and Mc-Math,² but they did not report the band origin.

The spectrometer used was a 15,000-line-per-inch grating spectrometer with a lead-sulfide detector. This instrument, previously described,3 was capable of resolving to better than 0.1 cm⁻¹ in the 2.2-µ (CS₂) region and to better than 0.2 cm^{-1} in the $1.4-\mu$ (CO₂) region.

A multiple-reflection absorption cell was used to obtain a long path length. In designing the cell a major consideration was to avoid the loss of aperture when small mirrors were used. The cell was designed for use in the spectral region beyond the photographic. This prevents the use of a long exposure time to compensate for any loss of aperture. In order to keep slit widths at a minimum and resolution at a maximum, the path length was kept to 6 m. With this path length it was possible to design a cell with negligible loss of aperture and still use small-diameter optics. A cell of long, variable pathlength and somewhat different optics has been described by White.4

Figure 1 is a schematic drawing of the cell with a central ray traced through the system. The lines representing this ray are numbered according to the number of the transversal of the cell they represent. The source, S, is placed off-axis and farther from the

spherical mirror, M_1 , than the radius, R, of this mirror. The radius of curvature of this mirror is 102 cm, and it has a diameter of 18 cm. Light from the source follows line 1 to M_1 , line 2 to mirrors M_2 and M_3 , line 3 back to M_1 , etc. The plane mirrors, M_2 , M_3 , M_4 , M_5 , and M_6 , are so arranged as to cause the final image of the source to be focussed on the entrance slit of the spectrometer. These plane mirrors were adjusted with the aid of a photocell, assuring maximum transmission of energy. Points X and X_2 are the points where the image of the source is focussed the first and second times. M_5 is placed considerably off-axis so that M_b can reflect the light out of the cell without intercepting light going to M_1 or M_3 . At no time is the off-axis condition as great as 8°. An aluminum tube 1 m long and 20 cm inside diameter forms the body of the cell.

An original record obtained on the recorder is shown in figure 2. The water absorption bands in this region distorted the CO₂ lines, and thus the various constants could not be calculated with the utmost accuracy. Table 1 shows the wavelength and vacuum wave number of the lines of this spec-

Table 1. Wavelengths and wave numbers of the 3v3 band of CO2

	R branch		P branch	
J	Wave- length	Wave number	Wave- length	Wave
	A	cm-1	A	cm^{-1}
0	14336. 3	6973. 39		
2	14333. 4	6974. 80		
4	14330. 5	6976. 21	14344.6	6969. 35
6	14327. 8	6977. 53	14348.4	6967. 51
8	14325. 0	6978. 89	14351.9	6965. 81
10	14322. 6	6980.06	14355.7	6963. 97
12	14320. 4	6981.13	14359.5	6962. 12
14	14318. 2	6982. 20	14363.8	6960.04
16	14316. 2	6983. 18	14368.1	6957. 69
18	14314. 6	6983. 96	14372.6	6955. 78
20	14312. 9	6984. 79	14377. 2	6953. 55
22	14311.5	6985. 47	14382.0	6951. 23
24	14310. 1	6986. 16	14387.0	6948. 82
26	14308. 9	6986.74	14392.3	6946. 26
28	14307. 9	6987. 23	14397. 5	6943. 75
30	14307. 0	6987. 67	14403.0	6941. 10
32	14306. 4	6987. 96	14408.4	6938. 49
34	*		14414.1	6935. 75
36			14420.0	6932. 91

E. F. Barker and T. Y. Wu, Phys. Rev. 45, 1 (1934).
 L. Goldberg, O. Mohler, A. Pierce, and R. McMath, Phys. Rev. 78, 74 (1950).
 Earle K. Plyler and Norman Gailar, J. Research NBS 47, 248 (1951) RP2249.
 John U. White, J. Opt. Soc. Am. 32, 285 (1942).

In analyzing the structure of the $3\nu_3$ band of CO_2 , use was made of the well-known combination differences for linear molecules 5

$$R(J-1)-P(J+1)=4B^{\prime\prime}(J+1/2),$$
 (1)

$$R(J)+P(J)=2\nu_0+2B'+2(B'-B'')J(J+1).$$
 (2)

Here B' is the upper and B'' the lower-state rotational constant, ν_0 the band origin, J the rotational quantum number of the lower state of the transition, and P and R are the short and long wave-number branches, respectively. These relations yield ν_0 6972.4 cm^{-1} , $\alpha_3 = 0.0032 \text{ cm}^{-1}$, $B_0 = 0.3900 \text{ cm}^{-1}$. These values are in good agreement with those reported by Benedict, Herman, and Silverman.6 Using the value of 2349.3 cm⁻¹ for ν_3 as reported by Herzberg (footnote 5), $X_{33} = -12.6$ cm⁻¹. The value

of 22.5 cm⁻¹ for X_{33} given by Benedict, Herman, and Silverman (footnote 6) is a typographical error and was intended to read 12.5 cm⁻¹.

The $3\nu_3$ band of CS₂ is presented in figure 3. Here five different band heads can be seen. The major

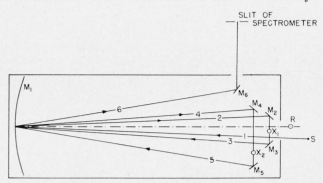
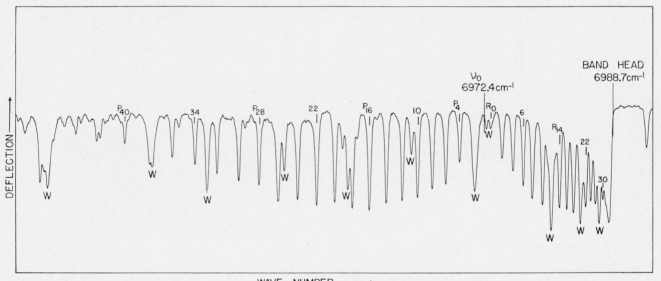


FIGURE 1. Schematic drawing of multiple reflection absorption cell.

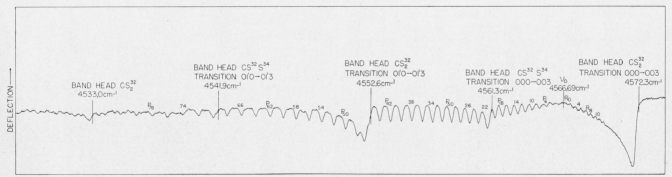
The lettered components are described in the text.



WAVE NUMBER

FIGURE 2. The 3v₃ absorption band of CO₂.

Some lines are labeled with the J values of the transition. Wavelengths and vacuum wave numbers of all the lines are found in table 1. Lines labeled Ware water absorption lines.



WAVE NUMBER-

FIGURE 3. The 3v3 absorption band of CS2.

Some lines are labeled with the J values of the transition. Wavelengths and vacuum wave numbers of all the lines are found in table 2.

⁵ G. Herzberg, Infrared and Raman spectra of polyatomic molecules, chapters III, IV (D. Van Nostrand Co., New York, N. Y., 1945).

⁶ W. S. Benedict, Robert C. Herman, and Shirleigh Silverman, J. Chem. Phys. 19,1325 (1951).

band head and the rotational structure are of the $000\rightarrow003$ transition and occur at about $572.3~\rm cm^{-1}$. At $4561.3~\rm cm^{-1}$ is the band head for the same transition for the isotopic molecule $CS^{32}S^{34}$. The band head of the transition $01^10\rightarrow01^13$ occurs at $4552.6~\rm cm^{-1}$, and the head for the same transition for the isotopic molecule occurs at $454~1.9~\rm cm^{-1}$. A very weak band head can be seen at $4533.0~\rm cm^{-1}$. This can be attributed to the pair of transitions $02^00\rightarrow02^03$ and $02^20\rightarrow02^23$. The band head of the transition $100\rightarrow103$ is expected to occur at approximately this position and may be superimposed at $4533.0~\rm cm^{-1}$.

The scarcity of lines in the R branch of the principal transition prevented the use of the combination differences (eq 1 and 2). Instead, the

absorption lines were fitted to the equation

$$\nu = \nu_0 + (B' + B'') m + (B' - B'') m^2,$$
 (3)

(see footnote 5) in order to determine ν_0 , B' and B''. These symbols have the same meanings as in eq 1 and 2. m=J+1 for the R branch, and m=-J for the P branch. A least-square fit to this equation yielded $\nu_0=4566.69~{\rm cm}^{-1}$, $B_0=0.1089~{\rm cm}^{-1}$, and $\alpha_3=(B''-B')/3=0.007~{\rm cm}^{-1}$. Table 2 contains the J and m values, the wavelengths, and vacuum wave numbers of the lines of this spectrum, and the wave

numbers calculated from eq 3.

The location of the band origins can be calculated to a good first approximation by assuming that the differences, $\nu_{\text{band head}} - \nu_0$, remain constant. This is shown by considering eq 3. A band head is expected when $d\nu/dm=0$ or when m=-(B'+B'')/2 (B'-B''). Substituting this value of m into eq 3, $\nu_{\text{band head}} - \nu_0 = -(B'+B'')^2/4(B'-B'')$ for the difference between the band head and the band origin. Comparing this difference for two transitions $(\nu_{\text{band head}} - \nu_0)_{000\to 00S} - (\nu_{\text{band head}} - \nu_0)_{01^10\to 01^13} = -(B_{003} + B_{000})^2/4 (B_{003} - B_{000}) + (B_{01^13} + B_{01^10})^2/4 (B_{01^13} - B_{01^10})$. Expressing these 4B's in terms of the equation $B(\nu) = B_e = \sum_1 \alpha_i (\nu_i + (d_i)/2)$ see footnote 5) and noting that $B_e \gg \alpha_1$, to a good approximation, we find $(\nu_{\text{band head}} - \nu_0)_{000\to 003} - (\nu_{\text{band head}} - \nu_0)_{01^10\to 01^13} < B_e$.

Since B_{ℓ} is approximately 0.1 cm⁻¹, it is seen that the difference between band head and band origin is equal for the transitions $000 \rightarrow 003$ and $01^{1}0 \rightarrow 01^{1}3$

to within experimental error.

Using this relation, ν_0 for $01^10 \rightarrow 01^13$ is found to be 4546.7 cm⁻¹. Taking the value of ν_0 for $000 \rightarrow 001$ of 1532.5 cm⁻¹ as measured at the Bureau on a prism spectometer, it is found that $X_{33} = -5.2$ cm⁻¹ and $X_{23} = -3.3$ cm⁻¹.

Table 2. Wavelengths band wave numbers of the $3\nu_3$ band of CS_2

J	m -	Wave- length	Wave number observed	Wave number calculated
R18 16 14 12 10	19 17 15 13 11	A 21875. 73 21876. 91 21878. 37 21879. 78 21881. 47	cm-1 4570.02 4569.78 4569.48 4569.18 4568.83	cm-1 4570.04 4569.76 4569.46 4569.15 4568.82
$\begin{array}{c} R8 \\ 6 \\ 4 \\ 2 \end{array}$	9 7 5 3	21883. 11 21884. 81 21886. 69 21888. 47	4568. 48 4568. 13 4567. 74 4567. 36	4568. 47 4568. 10 4567. 72 4567. 32
$P4 \\ 6 \\ 8 \\ 10 \\ 12$	$ \begin{array}{r} -4 \\ -6 \\ -8 \\ -10 \\ -12 \end{array} $	21895, 98 21898, 16 21900, 60 21903, 10 21905, 58	4565. 79 4565. 35 4564. 84 4564. 32 4563. 80	4565. 80 4565. 32 4564. 83 4564. 33 4563. 81
P14 16 18 20 22	$ \begin{array}{r} -14 \\ -16 \\ -18 \\ -20 \\ -22 \end{array} $	21908. 19 21910. 88 21913. 71 21916. 46	4563. 26 4562. 70 4562. 10 4561. 54	4563. 27 4562. 71 4562. 14 4561. 55 4560. 95
P24 26 28 30 32	$ \begin{array}{r} -24 \\ -26 \\ -28 \\ -30 \\ -32 \end{array} $	21922. 46 21925. 40 21928. 65 21931. 89 21935. 16	4560. 28 4559. 60 4559. 00 4558. 32 4557. 64	4560. 33 4559. 69 4559. 02 4558. 36 4557. 68
P34 36 38 40 42	-34 -36 -38 -40 -42	21938. 52 21941. 98 21945. 56 21949. 21 21952. 84	4556. 95 4556. 23 4555. 48 4554. 73 4553. 98	4556, 97 4556, 25 4555, 52 4554, 76 4553, 99
P44 46 48 50 52	$ \begin{array}{r} -44 \\ -46 \\ -48 \\ -50 \\ -52 \end{array} $	21956. 63 21964. 16 21968. 24 21972. 30	4553. 19 4551. 63 4550. 78 4549. 94	4553. 21 4552. 41 4551. 59 4550. 75 4549. 90
P54 56 58 60 62	-54 -56 -58 -60 -62	21976. 44 21980. 76 21985. 11 21989. 50 21994. 11	4549. 07 4548. 20 4547. 30 4546. 38 4545. 43	4549. 03 4548. 14 4547. 24 4546. 32 4545. 39
P64 66 68 70 72	$ \begin{array}{r} -64 \\ -66 \\ -68 \\ -70 \\ -72 \end{array} $	21998, 66 22003, 44 22008, 69 22012, 96 22018, 38	4544. 49 4543. 50 4542. 42 4541. 54 4540. 53	4544, 44 4543, 47 4542, 49 4541, 49 4540, 47
P74 76 78 80 82	-74 -76 -78 -80 -82	22023. 48 22028. 26 22033. 47 22038. 88 22044. 36	4539. 46 4538. 38 4537. 36 4536. 20 4535. 07	4539. 44 4538. 39 4537. 32 4536. 24 4535. 14
P84	-84	22049.99	4533. 91	4534. 02

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